CLAIMS

- 1. (original) A method of detecting an epigenetic abnormality associated with a disease comprising, identifying, within a eukaryotic genome, a locus having a hypomethylated sequence specific for said disease and an endogenous multi-copy DNA element.
- 2. (original) The method of claim 1, wherein said step of identifying comprises separate steps of identifying said disease-specific hypomethylated sequence and identifying said endogenous multi-copy DNA element.
- 3. (original) The method of claim 2, wherein the steps may be performed in any order.
- 4. (original) The method of claim 1, wherein said disease-specific hypomethylated sequence and said endogenous multi-copy DNA element are within 10 kilobases of separation.
- 5. (original) The method of claim 1, wherein said endogenous multi-copy DNA element is a retroelement that is normally methylated.
- 6. (original) The method of claim 5, wherein said retroelement is selected from the group consisting of endogenous retroviral sequences (ERV), SINE sequences, Alu sequences, LINE sequences, and L1 sequences.
- 7. (original) A method of identifying a chromosomal region associated with a disease state comprising:
- identifying a locus, within DNA obtained from said diseased sample, that has a DNA sequence that is hypomethylated and an endogenous multi-copy DNA element, wherein the DNA sequence is methylated in a non-disease sample and wherein the chromosomal region consists of from about 1 to about 10 DNA coding sequences that are proximal to the identified locus.
- 8. (original) A method of identifying a DNA coding sequence having an epigenetically altered expression pattern that contributes to a disease in an organism comprising: identifying a locus, within DNA obtained from said diseased sample, that has a DNA sequence that is hypomethylated and an endogenous multi-copy DNA element, said DNA sequence being methylated in a non-disease sample; and comparing expression patterns of the DNA coding sequence that comprises, or that is located proximal to, said identified locus within said diseased sample and said non-diseased sample, to identify said DNA coding sequence having an epigenetically altered expression pattern.
- 9. (original) The method of claim 8, wherein said disease is selected from the group consisting of Huntingdon's disease, schizophrenia, and bipolar disorder.

10. (original) A method of diagnosing an epigenetic abnormality correlated with a disease comprising:

identifying a DNA sequence that is hypomethylated within a locus that has an endogenous multi-copy DNA element and is obtained from a diseased sample, said DNA sequence being methylated in a non-disease sample.

- 11. (currently amended) The method Method of detecting an epigenetic abnormality as defined in claim 1, wherein said disease is associated with a non-Mendelian disease, and said method further comprising:
- a) extraction of genomic DNA from a sample that exhibits characteristics of a non-Mendelian disease;
- b) digestion of said genomic DNA with a methylation-sensitive restriction enzyme to produce a pool of restricted DNA fragments;
- c) fractionation of said pool of restricted DNA fragments to obtain DNA fragments of a desired size;
- d) amplification of at least a segment of said DNA fragments of a desired size with primers that anneal to an endogenous DNA element to produce a PCR product;
 - e) cloning of said PCR product into a sequencing vector;
- f) sequence determination of said PCR product to obtain a sequence of said PCR product;
- g) comparing said sequence against a genomic database to assign a locus for said epigenetic abnormality associated with a non-Mendelian disease.
- 12. (original) The method of claim 11, wherein said non-Mendelian disease is selected from the group consisting of schizophrenia, bipolar disorder, cancer, and diabetes.
- 13. (original) The method of claim 11, wherein said sample that exhibits characteristics of a non-Mendelian disease is brain tissue.
- 14. (original) The method of claim 13, wherein said sample that exhibits characteristics of a non-Mendelian disease is selected from the group consisting of frontal cortex and prefrontal cortex.
- 15. (original) The method of claim 11, wherein said desired size is less than 10 kb.
- 16. (original) The method of claim 11, wherein said endogenous DNA element is a multi-copy DNA element.
- 17. (original) The method of claim 16, wherein said multi-copy DNA element is selected from the group consisting of endogenous retroviral sequence, LINE, SINE, L1, and Alu.
- 18. (original) The method of claim 11, wherein said methylation-sensitive restriction enzyme is selected from the group consisting of AatlI (GACGTC); Bsh1236I (CGCG); Bsh1285I (CGRYCG); BshTI (ACCGGT); Bsp68I (TCGCGA); Bsp119I (TTCGAA);

Bsp143II (RGCGCY); Bsu15I (ATCGAT); Cfr10I (RCCGGY); Cfr42I (CCGCGG); CpoI (CGGWCCG); Eco47III (AGCGCT); Eco52I (CGGCCG); Eco72I (CACGTG); Eco105I (TACGTA); Ehel (GGCGCC); Esp3I (CGTCTC); FspAI (RTGCGCAY); Hin1I (GRCGYC); Hin6I (GCGC); HpaII (CCGG); Kpn2I (TCCGGA); MluI (ACGCGT); NotI (GCGGCCGC); NsbI (TGCGCA); PauI (GCGCGC); PdiI (GCCGGC); Pf123II (CGTACG); Psp1406I (AACGTT); PvuI (CGATCG); SaII (GTCGAC); SmaI (CCCGGG); SmuI (CCCGC); TaiI (ACGT); and TauI (GCSGC).

- 19. (currently amended) The method Method of claim 11, wherein said epigentic abnormality comprises identifying a gene having an epigenetically altered expression pattern that contributes to a to the non-Mendelian disease in an organism, said method further comprising:
- a) extraction of genomic DNA from a sample that exhibits characteristics of a non-Mendelian disease;
- b) digestion of said genomic DNA with a methylation sensitive restriction enzyme to produce a pool of restricted DNA fragments;
- e) fractionation of said-pool of restricted DNA fragments to obtain DNA fragments of a desired size;
- d) amplification of at least a segment of said DNA fragments of a desired size with primers that anneal to an endogenous DNA element to produce a PCR product;
- e) cloning of said PCR product into a sequencing vector;
- f) sequence determination of said PCR product to obtain a sequence of said PCR product;
- g) comparing said sequence against a genomic database to assign a locus for said epigenetic abnormality associated with a non-Mendelian disease;
 - h) searching said database to identify a gene located proximal to said locus;
- i) comparing expression patterns of said gene located proximal to said locus within a test sample that exhibits characteristics of said non-Mendelian disease with expression patterns of a corresponding gene within a control sample to identify said gene having an epigenetically altered expression pattern.

20. (canceled):

- 21. (currently amended) <u>A method</u> Method is isolating a probe for detecting an epigenetic abnormality associated with a non-Mendelian disease in accordance with claim 11, said method comprising:
- a) extraction of genomic DNA from a sample that exhibits characteristics of a non-Mendelian disease;
- b) digestion of said genomic DNA with a methylation-sensitive restriction enzyme to produce a pool of restricted DNA fragments;
- c) fractionation of said pool of restricted DNA fragments to obtain DNA fragments of a desired size;
- d) amplification of at least a segment of said DNA fragments of a desired size with primers that anneal to an endogenous DNA element to produce a PCR product;
- e) using said PCR product as said probe to detect said epigenetic abnormality associated with a non-Mendelian disease in another sample.

22. (canceled)

- 23. (original) A method of detecting a disease associated with an epigenetic abnormality comprising, identifying, within a eukaryotic genome, a locus having a hypomethylated sequence specific for said disease and an endogenous multi-copy DNA element.
- 24. (original) A method of diagnosing a disease correlated with an epigenetic abnormality comprising:

identifying a DNA sequence that is hypomethylated within a locus that has an endogenous multi-copy DNA element and is obtained from a diseased sample, said DNA sequence being methylated in a non-disease sample.